

10/666, 068

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FILE 'HOME' ENTERED AT 13:19:57 ON 10 NOV 2004

=> fil reg  
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ENTRY SESSION

FULL ESTIMATED COST

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FILE 'REGISTRY' ENTERED AT 13:20:05 ON 10 NOV 2004  
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STRUCTURE FILE UPDATES: 8 NOV 2004 HIGHEST RN 777024-10-9  
DICTIONARY FILE UPDATES: 8 NOV 2004 HIGHEST RN 777024-10-9

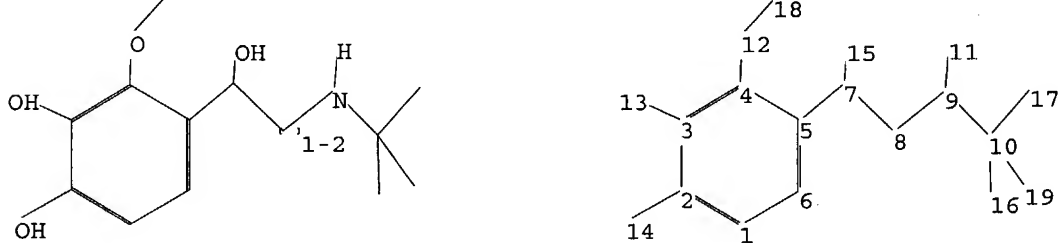
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<http://www.cas.org/ONLINE/DBSS/registryss.html>

=>  
Uploading C:\Program Files\Stnexp\Queries\10666068.str



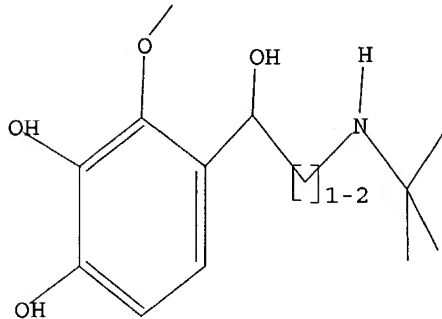
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7 8 9 10 11 12 13 14 15 16 17 18 19  
ring nodes :  
1 2 3 4 5 6  
chain bonds :  
2-14 3-13 4-12 5-7 7-8 7-15 8-9 9-10 9-11 10-16 10-17 10-18 12-18  
ring bonds :  
1-2 1-6 2-3 3-4 4-5 5-6  
exact/norm bonds :  
2-14 3-13 4-12 7-15 8-9 9-10 12-18  
exact bonds :  
5-7 7-8 9-11 10-16 10-17 10-19  
normalized bonds :  
1-2 1-6 2-3 3-4 4-5 5-6

Match level :  
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:CLASS  
11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS  
19:CLASS

L1 STRUCTURE UPLOADED

=> d query

L1 STR



Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 13:20:22 FILE 'REGISTRY'  
SAMPLE SCREEN SEARCH COMPLETED - 6 TO ITERATE

100.0% PROCESSED 6 ITERATIONS  
SEARCH TIME: 00.00.01

1 ANSWERS

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*  
BATCH \*\*COMPLETE\*\*  
PROJECTED ITERATIONS: 6 TO 266  
PROJECTED ANSWERS: 1 TO 80

L2 1 SEA SSS SAM L1

=> s 11 full  
FULL SEARCH INITIATED 13:20:27 FILE 'REGISTRY'  
FULL SCREEN SEARCH COMPLETED - 91 TO ITERATE

100.0% PROCESSED 91 ITERATIONS  
SEARCH TIME: 00.00.01

16 ANSWERS

L3 16 SEA SSS FUL L1

=> fil caplus  
COST IN U.S. DOLLARS SINCE FILE TOTAL  
ENTRY SESSION  
FULL ESTIMATED COST 155.42 155.63

FILE 'CAPLUS' ENTERED AT 13:20:30 ON 10 NOV 2004  
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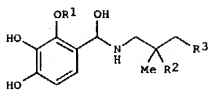
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FILE COVERS 1907 - 10 Nov 2004 VOL 141 ISS 20  
FILE LAST UPDATED: 9 Nov 2004 (20041109/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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=> s 13
L4          9 L3
=> d 14 1-9 abs ibib hitstr
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I

AB Title compds. [I]; R1, R2 = Cl-4 alkyl; R3 = Cl-4 alkyl, (substituted) Ph; or R2R3 = CH2CH2, (CH2)3, were prepared as  $\beta$ 2-adrenergic sympathomimetics (no data). Thus, 1-(3,4-dihydroxy-2-methoxyphenyl)-2-(1,1-dimethylpropylamino)ethanone (preparation given) was hydrogenated by using

PtO in MeOH to give 85% 4-[2-(1,1-dimethylpropylamino)-1-hydroxyethyl]-3-methoxybenzene-1,2-diol.

ACCESSION NUMBER: 2004:307317 CAPLUS

DOCUMENT NUMBER: 140:321101

TITLE: Preparation of benzenediols for treatment of respiratory tract diseases

INVENTOR(S): Bouysou, Thierry; Buettnner, Frank; Konetzki, Ingo; Pestel, Sabine; Schnapp, Andreas; Schollenberger, Hermann; Schromm, Kurt; Heine, Claudia

PATENT ASSIGNEE(S): Boehringer Ingelheim Pharma GmbH & Co. Kg, Germany  
Ger. Offen., 14 pp.

DOCUMENT TYPE: Patent

LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 10246374	A1	20040415	DE 2002-10246374	20021004
US 2004121208	A1	20040624	US 2003-666068	20030919
WO 2004033412	A1	20040422	WO 2003-EP10661	20030925
W: AE, AG, AL, AM, AT, AU, A2, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, N2, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, U2, VC, VN, YU, ZA, ZM, ZW, AM, A2, BY, KG, KZ, MD				
W: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, C2, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BE, BA, CF, CG, CI, CM, GA, GN, GO, GW, ML, MR, NE, SN, TD, TG				

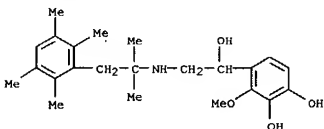
PRIORITY APPLN. INFO.: DE 2002-10246374 A 20021004  
US 2002-432499P P 20021211

OTHER SOURCE(S): MARPAT 140:321101

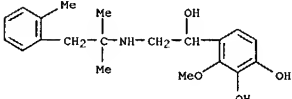
IT 677776-89-5P 677777-04-7P 677777-17-2P

677777-23-0P 677777-27-4P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

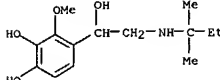


RN 677777-27-4 CAPLUS  
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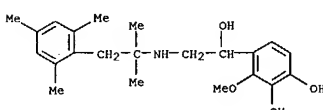


L4 ANSWER 1 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)  
(Uses)  
(prepn. of benzenediols for treatment of respiratory tract diseases)

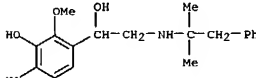
RN 677776-89-5 CAPLUS  
CN 1,2-Benzenediol, 4-[2-[(1,1-dimethylpropyl)amino]-1-hydroxyethyl]-3-methoxy- (9CI) (CA INDEX NAME)



RN 677777-04-7 CAPLUS  
CN 1,2-Benzenediol, 4-[2-[(1,1-dimethyl-2-(2,4,6-trimethylphenyl)ethyl]amino]-1-hydroxyethyl]-3-methoxy- (9CI) (CA INDEX NAME)

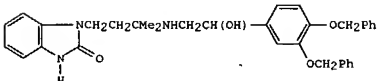
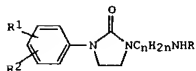


RN 677777-17-2 CAPLUS  
CN 1,2-Benzenediol, 4-[2-[(1,1-dimethyl-2-phenylethyl)amino]-1-hydroxyethyl]-3-methoxy- (9CI) (CA INDEX NAME)



RN 677777-23-0 CAPLUS  
CN 1,2-Benzenediol, 4-[2-[(1,1-dimethyl-2-(2,3,5,6-tetramethylphenyl)ethyl]amino]-1-hydroxyethyl]-3-methoxy- (9CI) (CA INDEX NAME)

GI



AB The title compds. I [R = (un)substituted  $\beta$ -hydroxyphenethyl, R1 = H, halo, Cl-4 alkyl or alkoxy, F3C, NH2, R2 = H, halo, Cl-4 alkyl or alkoxy, F3C, R1R2 = methylenedioxy, ethylenedioxy, n = 2-6] useful as antihypertensives, broncholytics, and vasodilators (no data) were prepared

by a variety of reduction reactions. Thus, in a comparative example 3,4-(PhCH2O)2C6H3COCH(OH)OBt was heated with 1-(3-amino-3-methylbutyl)benzimidazolidinone in EtOH 3 h, cooled, and treated with NaBH4 to give II, isolated as its maleate.

ACCESSION NUMBER: 1983:488195 CAPLUS

DOCUMENT NUMBER: 99:88195

TITLE: N-Aminomethylimidazolidines

INVENTOR(S): Mentrup, Anton; Schromm, Kurt; Renth, Ernst Otto; Reichl, Richard; Trauecker, Werner; Hoefke, Wolfgang

PATENT ASSIGNEE(S): Boehringer Ingelheim International G.m.b.H., Fed. Rep.

Ger. Pat. Specif. (Aust.), 66 pp.

CODEN: ALXXAP

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
AU 526579	B2	19830120	AU 1981-67647	19810225
AU 8167647	A1	19810521		

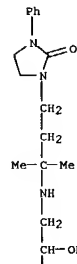
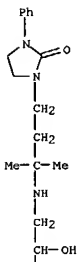
PRIORITY APPLN. INFO.: AU 1981-67647 19810225

IT 64928-21-8P 86733-03-1P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 64928-21-8 CAPLUS

CN 2-Imidazolidinone, 1-[3-[(2-(3,4-dihydroxy-2-methoxyphenyl)-2-hydroxyethyl)amino]-3-methylbutyl]-3-phenyl- (9CI) (CA INDEX NAME)



PAGE 2-A



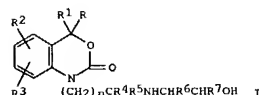
PAGE 2-A



RN 86733-03-1 CAPLUS  
CN Formic acid, compd. with 1-[3-((2-(3,4-dihydroxy-2-methoxyphenyl)-2-hydroxyethyl)amino)-3-methylbutyl]-3-phenyl-2-imidazolidinone (9CI) (CA INDEX NAME)  
CM 1  
CRN 64-18-6  
CMF C23 H31 N3 O5

CM 2  
CRN 64-18-6  
CMF C H2 O2

O=CH-OH



AB Benzoxazinones I (R, R1, R6 = H, alkyl; R2, R3 = H, F, Cl, OH, Me, Et, alkoxy; R2R3 = OCH2O; R4, R5 = H, Me; R7 = substituted Ph; n = 1-3) were prepared. Thus 1,1-dimethyl-3-(4,4-dimethyl-2-oxo-3,1-benzoxazin-1-yl)propanamine was treated with 3,4-H2NCO(HO)C6H3COCH2Br and reduced with NaBH4 to give I [R = R1 = R5 = Me, R2 = R3 = R6 = H, R7 = 3,4-H2NCO(HO)C6H3, n = 2] (II). II·MeSO3H had antihypertensive activity.

at 10 mg/kg orally in rats.

ACCESSION NUMBER: 1982:199711 CAPLUS  
DOCUMENT NUMBER: 96:199711  
TITLE: 3,1-Benzoxazin-2-ones and their uses  
INVENTOR(S): Mentrup, Anton; Schromm, Kurt; Renth, Ernst Otto; Hoefke, Wolfgang; Gaida, Wolfram; Streller, Ilse; Fuegner, Armin  
PATENT ASSIGNEE(S): Boehringer, C. H., Sohn, Fed. Rep. Ger.  
SOURCE: Eur. Pat. Appl.  
DOCUMENT TYPE: Patent  
LANGUAGE: German  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

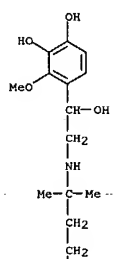
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 43940	A1	19820120	EP 1981-104787	19810622
EP 43940	B1	19840912		
R: AT, BE, CH, DE, FR, IT, LU, NL, SE				
DE 3026534	A1	19820318	DE 1980-3026534	19800712
AT 9336	E	19840915	AT 1981-104787	19810622
US 4341778	A	19820727	US 1981-280349	19810706
DK 8103067	A	19820113	DK 1981-3067	19810710
DK 149851	B	19861013		
DK 149851	C	19870504		
FI 8102183	A	19820113	FI 1981-2183	19810710
FI 74703	B	19871130		
FI 74703	C	19880310		
NO 8102355	A	19820113	NO 1981-2355	19810710
NO 158578	B	19880627		
NO 158578	C	19881005		
GB 2080296	A	19820203	GB 1981-21321	19810710
GB 2080296	B2	19830928		
ES 503837	A1	19820601	ES 1981-503837	19810710
AU 8172731	A1	19820916	AU 1981-72731	19810710
AU 540916	B2	19841206		
ZA 8104687	A	19830330	ZA 1981-4687	19810710
DD 202018	A5	19830824	DD 1981-231670	19810710

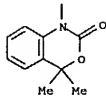
L4 ANSWER 3 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)  
HU 25946 O 19830829 HU 1981-2036 19810710  
HU 183515 B 19840528  
CA 1165317 A1 19840410 CA 1981-381559 19810710  
IL 63285 A1 19850331 IL 1981-63285 19810710  
JP 57048975 A2 19820320 JP 1981-109186 19810713  
ES 508653 A1 19821101 ES 1982-508653 19820112  
ES 508654 A1 19821101 ES 1982-508654 19820112  
ES 508655 A1 19821101 ES 1982-508655 19820112  
DE 1980-3026534 D 19800712 19800712  
EP 1981-104787 EP 1981-104787 19810622

OTHER SOURCE(S): CASREACT 96:199711  
IT 81696-95-9P  
RL: SPN (Synthetic preparation); PREP (Preparation)  
(preparation of)  
RN 81696-95-9 CAPLUS  
CN Formic acid, compd. with 1-[3-((2-(3,4-dihydroxy-2-methoxyphenyl)-2-hydroxyethyl)amino)-3-methylbutyl]-3-phenyl-2-imidazolidinone (1:1) (9CI) (CA INDEX NAME)

CM 1  
CRN 81696-94-8  
CMF C24 H32 N2 O6

PAGE 1-A

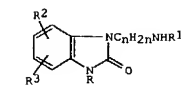




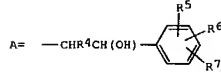
CM 2

CRN 64-18-6  
CMF C H2 O2

O=CH-OH



I



AB Title compds. I [R = H, alkyl; n = 2-6; R1 = H, PhCH2, A (R4 = H, Me, Et, R5, R6, and R7 (same or different) are H, halo, CH2OH, CF3, alkyl, alkoxy, NO2, cyano, CONHR8 (R8 = H, alkyl, OH), CO2H, carbalkoxy, OH, alkaneoxy, alkoxy, PhCH2O, MeSO2CH2; or R5R6 = OCH2O, OCH2CH2O, benzo, OCH2CONH, CH2CH2CONH]; R2 = H, halo, alkyl, alkoxy, CF3, NH2; R3 = H, halo, alkyl, alkoxy, CF3; or R2R3 = OCH2O, OCH2CH2O], useful as central nervous system stimulants, antihypertensives, and vasodilators (no data), were prepared by different methods. A mixt of 3,4-(PhCH2O)2C6H3COCH(OH)OEt and 1-(3-amino-3,3-dimethylpropyl)-2-benzimidazolinone in EtOH was heated 3 h, mixed with NaBH4 at 0-5°, kept 12 h at room temperature, acidified, and worked up to give I [R = R2 = R3 = H, CnH2n = CH2CH2CMe2, R1 = A (R4 = R7 = H, R5 = 3-PhCH2O, R6 = 4-PhCH2O)]. Also prepared was I [R = R2 = R3 = H, CnH2n = CH2CH2CMe2, R1 = A (R4 = R7 = H, R5 = 3-OH, R6 = 4-OH)], which exhibited bronchodilator activity.

ACCESSION NUMBER: 1981:4017 CAPLUS  
DOCUMENT NUMBER: 94:4017  
TITLE: Aminoalkyl-substituted benzimidazolidin-2-ones  
INVENTOR(S): Hoecke, Wolfgang; Mentrup, Anton; Reichl, Richard; Renth, Ernst Otto; Schromm, Kurt; Trauneker, Werner  
PATENT ASSIGNEE(S): Boehringer Ingelheim G.m.b.H., Fed. Rep. Ger.  
SOURCE: U.S., 45 pp. Cont.-in-part of U.S. 4,154,829.  
CODEN: USXXAM  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 2  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4215119	A	19800729	US 1979-26608	19790403
DE 2609645	A1	19770915	DE 1976-2609645	19760309
US 4154829	A	19790515	US 1977-773394	19770302
US 4271158	A	19810602	US 1979-102904	19791213
US 4363814	A	19821214	US 1980-218786	19801222
PRIORITY APPLN. INFO.:				DE 1976-2609645 19760309

US 1979-26608 19790403  
US 1979-102904 19791213

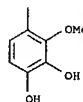
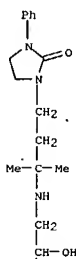
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CRN 64-18-6  
CMF C H2 O2

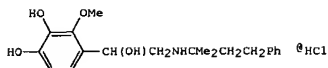
IT 64928-22-9P  
RL: SPN (Synthetic preparation); PREP (Preparation)  
(preparation of)  
RN 64928-22-9 CAPLUS  
CN Formic acid, compd. with 1-[3-((2-(3,4-dihydroxy-2-methoxyphenyl)-2-hydroxyethyl)amino)-3-methylbutyl]-3-phenyl-2-imidazolidinone (1:1) (9CI)  
(CA INDEX NAME)

CM 1

CRN 64928-21-8  
CMF C23 H31 N3 O5

O=CH-OH





AB The  $\beta$ -receptor stimulant effects of Sm220Cl-HCl (dl-N-(1,1-dimethyl-3-phenyl-propyl)-2-hydroxy-2-(3,4-dihydroxy-2-methoxyphenyl)ethylamine-HCl) (I) [64725-05-9] and (-)-isoprenaline were compared in isolated atrial (H1) and tracheal (H2) preps. from guinea-pigs and cats. The compds. were also tested for their ability to increase the heart rate (H1), reduce serotonin-induced increases in pulmonary resistance (H2), and decrease soleus muscle contractility (H2) in vivo in the two species. Calculated selectivity ratios

[activity-ratio (heart):activity-ratio (bronchial smooth muscle)] from the in vitro expts.

showed that I possessed  $\beta$ 2-receptor selectivity. This was more marked in guinea pig than in cat preps. In the anesthetized animals

this species difference was more apparent; in cats, I was non-selective in its actions for  $\beta$ 1- and  $\beta$ 2-receptor mediated responses, while marked  $\beta$ 2-receptor selectivity was obtained in the guinea pig. Since in both species, the activity-ratios for  $\beta$ 2-receptor mediated actions are similar, the differences in the  $\beta$ 1/ $\beta$ 2-receptor selectivity of (I) are caused by the divergent cardiac effects produced by the drug.

ACCESSION NUMBER: 1978:310 CAPLUS

DOCUMENT NUMBER: 88:310

TITLE: Species difference in the  $\beta$ 1/ $\beta$ 2-adrenoceptor

selectivity of Sm220Cl in the cat and guinea-pig

AUTHOR(S): Bohmer, K.; Raper, C.

CORPORATE SOURCE: Dep. Pharmacol., Victorian Coll. Pharm., Parkville,

Australia

SOURCE: Clinical and Experimental Pharmacology and Physiology

(1977), 4(4), 349-58

CODEN: CEXPB9; ISSN: 0305-1870

DOCUMENT TYPE: Journal

LANGUAGE: English

IT 24008-01-3

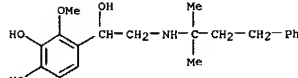
RL: BIOL (Biological study)  
(sympathomimetic activity of, selectivity of, species differences in)

RN 24008-01-3 CAPLUS

CN 1,2-Benzenediol,

4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-

3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

AB Approx. 270 title compds., RCH(OH)CH2NH2R1 (I, R = aryl, e.g., p-HOC6H4, 3,5-(PhCH2)2C6H3, 3,4-C12C6H3; n = 1-3, CH2CH2CMe2, etc.; R1 = 1,2,3,4-tetrahydro-2-oxquinolino (Q), 2-oxo-1,2-dihydrobenzimidazol-1-yl, 2-oxo-3-phenylimidazolin-1-yl, etc.) were prepared from R1ZNH2 and RCOCHO or its derivs., or RCOCH2Br. Thus, 5.6 g

3,4-dichlorophenylglyoxal

hydrate and 4.5 g 1-(3-aminopropyl)-1,2,3,4-tetrahydro-2-quinolinone was heated 1 hr at 50° and the mixture treated with 5 g NaBH4 at 0° to give 5.9 g I (R = 3,4-C12C6H3, Z = (CH2)n, n = 1-3, R1 = Q). I (R = 3,4-MeSO2NH(OH)C6H3, Z = CH2CH2CMe2, R1 = Q) was 18X as effective as Isosuprime as a peripheral vasodilator in the dog. I (R = 3,4-MeNHCO(OH)C6H3, Z = CH2CH2CMe2, R1 = Q) produced a blood pressure of 85 mm when given to hypertensive rats. Guinea pigs treated with I (R = 3,4-(HO)C6H3, Z = CH2CH2CMe2, R1 = 2,3-dihydro-2-oxo-benzimidazol-1-yl), exhibited a broncholytic ED50 (intravenous) of 0.09  $\mu$ g/kg vs. 3  $\mu$ g/kg for Isoproterenol.

ACCESSION NUMBER: 1977:601503 CAPLUS

DOCUMENT NUMBER: 87:201503

TITLE: Aminoalkyl heterocycles

INVENTOR(S): Mentrup, Anton; Schromm, Kurt; Renth, Ernst Otto;

Reichl, Richard; Trauecker, Werner; Hoefke, Wolfgang

PATENT ASSIGNEE(S): Boehringer, C. H., Sohn, Fed. Rep. Ger.

SOURCE: Ger. Offen., 79 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent

LANGUAGE: German

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2609645	A1	19770915	DE 1976-2609645	19760309
SU 698530	D	19791115	SU 1977-2453505	19770222
FI 7700566	A	19770910	FI 1977-586	19770223
FI 69709	B	19950830		
FI 69707	C	19951210		
AT 7701223	A	19900615	AT 1977-1223	19770224
AT 360542	B	19910112		
RO 76589	P	19810430	RO 1977-96539	19770301
RO 70569	P	19811124	RO 1977-89565	19770301
RO 79706	P	19820817	RO 1977-101591	19770301
US 4154829	A	19790515	US 1977-773394	19770302
CS 209435	P	19811231	CS 1977-1476	19770304
CS 220320	P	19830325	CS 1978-5352	19770304
NL 7702403	A	19770913	NL 1977-2403	19770307
CH 630358	A	19820615	CH 1977-2819	19770307
BE 852223	A1	19770908	BE 1977-175593	19770308
DK 7701021	A	19770910	DK 1977-1021	19770308
JP 52108970	A2	19770912	JP 1977-25323	19770308
NO 7700804	A	19770912	NO 1977-804	19770308
NO 147950	B	19830405		
NO 147950	C	19830713		
AU 7723009	A1	19780914	AU 1977-23009	19770308
AU 515953	B2	19810514		
ZA 7701412	A	19781129	ZA 1977-1412	19770308
CA 1086317	A1	19800923	CA 1977-273388	19770308
IL 51627	A1	19801026	IL 1977-51627	19770308
PL 112937	B1	19801129	PL 1977-215210	19770308
HU 20328	O	19810728	HU 1977-B01653	19770308

HU 177953 P 19820228

SE 435059 B 19840903 SE 1977-2609 19770308

SE 435059 C 19841213

FR 2343731 A1 19771007 FR 1977-7018 19770309

FR 2343731 B1 19820226

GB 1571231 A 19800709 GB 1977-9952 19770309

SU 676163 D 19790725 SU 1977-2541704 19771116

SU 683616 D 19790830 SU 1977-2543651 19771116

SU 685149 D 19790905 SU 1977-2542149 19771116

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FR 2372810 B1 19821126

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ES 466602 A1 19781001 ES 1978-466606 19780203

US 4215119 A 19800729 US 1979-26608 19790403

US 4271578 A 19810602 US 1979-102904 19791213

AT 8000203 A 19810215 AT 1980-203 19800116

AT 363940 B 19810910

AT 8000204 A 19810215 AT 1980-204 19800116

AT 363941 B 19810910

AT 8000207 A 19810215 AT 1980-207 19800116

AT 363942 B 19810910

AT 8000208 A 19810215 AT 1980-208 19800116

AT 363943 B 19810910

AT 8000209 A 19810215 AT 1980-209 19800116

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AT 372084 A 19830115 AT 1980-206 19800116

US 4363814 A 19821214 US 1980-218786 19801222

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CH 630360 A 19820615 CH 1981-2847 19810430

CH 630361 A 19820615 CH 1981-2848 19810430

CH 630362 A 19820615 CH 1981-2849 19810430

CH 630363 A 19820615 CH 1981-2850 19810430

CH 630364 A 19820615 CH 1981-2851 19810430

CH 630365 A 19820615 CH 1981-2852 19810430

JP 61000072 A2 19860106 JP 1985-126401 19850612

DE 1976-2609645 DE 1976-2609645 19760309

AT 1977-1223 19770224

US 1977-773394 19770302

CH 1977-2819 19770307

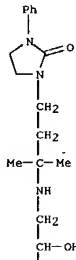
US 1979-26608 19790403

US 1979-102904 19791213

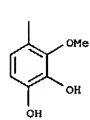
IT 64920-22-98  
RL: SPN (Synthetic preparation); PREP (Preparation)

L4 ANSWER 6 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)  
 RN (prep. of)  
 64928-22-9 CAPLUS  
 CN Formic acid, compd. with 1-[{2-(3,4-dihydroxy-2-methoxyphenyl)-2-hydroxyethyl]amino)-3-methylbutyl]-3-phenyl-2-imidazolidinone (1:1) (9CI)  
 (CA INDEX NAME)  
 CM 1  
 CRN 64928-21-8  
 CMF C23 H31 N3 O5

PAGE 1-A



L4 ANSWER 6 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)  
 O=CH-OH



PAGE 2-A

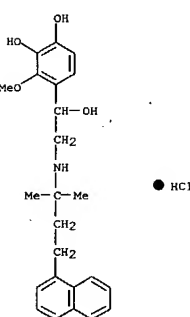
CM 2  
 CRN 64-18-6  
 CMF C H2 O2

L4 ANSWER 7 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN  
 GI For diagram(s), see printed CA issue.  
 AB The title compds. (I) were prepared by catalytic hydrogenation of the corresponding amino ketones. Thus,  $\alpha$ -bromo-4-(benzoyl)acetophenone and N-[2-(1-naphthyl)-ethyl]benzylamine was refluxed in MeCN and the product hydrogenated (Pd-C) to give I [Q = (CH2)2, R = R1 = R2 = H]. Similarly prepared were 23 other I and 2-  
 naphthyl analogs.  
 ACCESSION NUMBER: 1971:488382 CAPLUS  
 DOCUMENT NUMBER: 75:88382  
 TITLE: Pharmacologically active naphthylalkylamines  
 INVENTOR(S): Schiromm, Kurt; Mentrup, Anton; Renth, Ernst O.; Trauecker, Werner  
 PATENT ASSIGNEE(S): Boehringer, C. H., Sohn  
 SOURCE: Ger. Offen., 23 pp.  
 CODEN: GWXXBX  
 DOCUMENT TYPE: Patent  
 LANGUAGE: German  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 1962497	A	19710616	DE 1969-1962497	19691212
DE 1962497	C3	19790920		
DE 1962497	B2	19790125		
CH 556322	A	19741129	CH 1970-18189	19701209
CH 556323	A	19741129	CH 1973-4776	19701209
CH 564509	A	19750731	CH 1973-4777	19701209
SE 378101	B	19750818	SE 1970-16667	19701209
NL 7018031	A	19710615	NL 1970-18031	19701210
NL 169583	B	19820301		
NL 169583	C	19820802		
HU 162736	P	19730428	HU 1970-BO1262	19701210
RO 56908	P	19750315	RO 1970-65259	19701210
RO 61132	P	19761215	RO 1970-68643	19701210
RO 61063	P	19780715	RO 1970-68644	19701210
FR 2081347	A1	19711203	FR 1970-44709	19701211
FR 2081347	A5	19711203		
AT 299924	B	19720710	AT 1970-11175	19701211
AT 302284	B	19721010	AT 1971-7400	19701211
SU 384229	D	19730523	SU 1970-1497851	19701211
GB 1330188	A	19730912	GB 1970-59091	19701211
CS 151062	P	19730917	CS 1970-8374	19701211
CS 151063	P	19730917	CS 1971-7139	19701211
CS 151064	P	19730917	CS 1971-7140	19701211
ES 386345	A1	19740101	ES 1970-35645	19701211
SU 419326	D	19740229	SU 1970-1739938	19701211
IL 358400	A1	19740630	IL 1970-35840	19701211
AT 317192	B	19740812	AT 1971-7401	19701211
NO 131126	B	19741230	NO 1970-4790	19701211
PL 81424	P	19750830	PL 1970-144934	19701211
SU 055449	D	19760228	SU 1970-1239929	19701211
PL 84354	P	19760331	PL 1970-174664	19701211
PL 84355	P	19760331	PL 1970-174663	19701211
JP 51016420	B4	19760524	JP 1970-110358	19701211
DK 136526	B	19771024	DK 1970-6320	19701211
FI 53301	B	19771230	FI 1970-3343	19701211
FI 53301	C	19780410		
ES 395482	A1	19731216	ES 1971-395482	19710928

L4 ANSWER 7 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)  
 ES 395481 A1 19731216 ES 1971-395481 19710928  
 JP 51038716 B4 19761023 JP 1973-59198 19730525  
 JP 52021505 B4 19770610 JP 1973-59199 19730525  
 US 3966814 A 19760629 US 1973-373933 19730627  
 PRIORITY APPLN. INFO.: DE 1969-1962497 19691212  
 US 1970-92527 19701124

IT 33457-03-3P RL: SBN (Synthetic preparation); PREP (Preparation)  
 . (preparation of)  
 RN 33457-03-3 CAPLUS  
 CN Benzyl alcohol,  $\alpha$ -{[(1,1-dimethyl-3-(1-naphthyl)propyl]amino)methyl]-3,4-dihydroxy-2-methoxy-, hydrochloride (8CI) (CA INDEX NAME)-



L4 ANSWER 8 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN  
 GI For diagram(s), see printed CA Issue.  
 AB The title compds. (I) were prepared (1) by the reaction of QCOCHR3X (R = R1 or a protective group; X = halogen) and HNR'R4 (II) (R' = H or benzyl), followed by reduction; or (2) by reduction of QR5COR3 (R5 = CO or CHO) and R4NH2 or of the Schiff base condensed from both; or (3) by reaction of QE (E = 2-R3-substituted-1,2-epoxyethyl or CHOCHCR3) and II, followed by removal of the protective groups; or (4) by the reaction under reduction of QCH(OH)CHR3NH2 with R6COR7 (R6 = H or straight-chain lower alkyl, R7 = lower alkyl or 1,4-benzodioxan-2-yl); or (5) by reduction of QR5CONHR4, when the protective group RO = acetal or benzyl ether, removal of the protective groups; or (6) by the reaction of QCH(OH)CHR3.NHR' with R4Y (Y = halo or an acid radical) in the presence of excess amine, Na2CO3, or K2CO3 and elimination of the protective groups. Thus, the Na salt of 2-hydroxy-3,4-diphenylmethoxyacetophenone was reacted with EtOH and EtI to give 2-ethoxy-3,4-diphenylmethylenedioxyacetophenone (III), m. 82°. III (72 g) was reacted at 60° with 10 ml Br and then 60 g PhCH2NH-Pr-iso to yield

1-(2-ethoxy-3,4-diphenylmethylenedioxy)phenyl 1)-1-oxo-2-(benzylisopropylamino)ethane (IV). After purifying 81 g IV in 540 ml MeOH and 270 ml H2O over animal C, the solution was hydrogenated over

Pd/C, to yield 1-(2-ethoxy-3,4-dihydroxyphenyl)-1-oxo-2-(isopropylamino)ethane (V), hydrochloride m. 203-5° (95% iso-PrOH). V (20 g) was hydrogenated over Pt to yield 18 g 1-(2-ethoxy-3,4-dihydroxyphenyl)-1-hydroxy-2-(isopropylamino)ethane (I, R1 = R3 = H, R2 = EtO, R4 = iso-Pr), hydrochloride m. 184° (EtOH). Similarly prepared were the following I (R1, R2, R3, R4, and m.p. of hydrochloride, unless mentioned otherwise, given): H, Pr, H, iso-Pr, 158-9°; H, Me, H, 3-phenylpropyl, 176°; H, Me, H, 2-(hydroxyphenyl)isopropyl, (benzoate) 110°; H, MeO, H, cyclopentyl, 161-2°; H, MeO, H, phenoxethyl, 87-8° (crystallized with 0.5 mole Me2CO); H, MeO, H, tert-Bu, 97-9° (base) (0.5 mole H2O of crystallization); H, MeO, H, p-tolylxyethyl, 109-11°; H, MeO, H, o-tolylxyethyl, 134-5°; H, MeO, H, m-tolylxyethyl, 126-7°; H, MeO, H, o-methoxyphenoxethyl, 78-80° (crystallized with 1 mol MeCN); H, MeO, H, 1,1-dimethyl-3-phenylpropyl, 175-6°; H, MeO, H, 1,1-dimethyl-3-p-tolylpropyl, 168-70°; H, Me, H, tert-Bu, (benzoate) 179-81°; H, MeO, Et, iso-Pr, 220-2°; Ac, Me, H, iso-Pr, 99°; Ac, MeO, H, iso-Pr. Prepared intermediates are (m.p. or b.p. given): 3-allyloxy-4-methoxyacetophenone, bp 180-2°; 2-propyl-3-hydroxy-4-methoxyacetophenone, 87-9°; 2-propyl-3-hydroxy-4-methoxyacetophenone, ; 2-propyl-3-acetoxy-4-methoxybromoacetophenone, ; 1-(2-propyl-3-acetoxy-4-methoxyphenyl)-1-oxo-2-(benzylisopropylamino)ethane, ; 1-(2-propyl-3-hydroxy-4-methoxyphenyl)-1-oxo-2-(benzylisopropylamino)ethane-HCl, 100°; 1-(2-propyl-3-hydroxy-4-methoxyphenyl)-1-oxo-2-(isopropylamino)ethane-HCl, 93-5°; 1-(2-propyl-3,4-dihydroxyphenyl)-1-oxo-2-(isopropylamino)ethane-HCl, 181-2°;  $\alpha$ -bromo-2-methyl-3,4-dimethoxyacetophenone,  $\alpha$ -(benzyl-3-methyl-3,4-dimethoxyacetophenone)-(HO2C)2, 118-21°;  $\alpha$ -(3-phenylpropylamino)-2-methyl-3,4-dimethoxyacetophenone-HCl, 210-17°;  $\alpha$ -(3-phenylpropylamino)-2-methyl-3,4-dihydroxyacetophenone-HBr, 179° (base m. 130-8°);

L4 ANSWER 8 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)  
 CH 507200 A 19710515 CH 1967-507200 19671016  
 CH 523219 A 19720531 CH 1967-523219 19671016  
 CH 548365 A 19740430 CH 1972-4528 19671016  
 DK 130070 B 19741216 DK 1967-5161 19671017  
 BE 705312 A 19680419 BE 1967-705312 19671018  
 NL 6714161 A 19680419 NL 1967-14161 19671018  
 NL 158480 B 19781115 19671018  
 GB 1204195 A 19700903 GB 1967-1204195 19671018  
 SE 368196 B 19740624 SE 1967-14280 19671018  
 SE 380792 B 19751117 SE 1971-6612 19671018  
 ES 360964 A1 19701016 ES 1968-360964 19681130  
 ES 360961 A1 19701116 ES 1968-360961 19681130  
 ES 360962 A1 19701116 ES 1968-360962 19681130  
 ES 360963 A1 19701116 ES 1968-360963 19681130  
 ES 360965 A1 19701116 ES 1968-360965 19681130  
 PRIORITY APPLN. INFO.: DE 1966-B89417 A 19661018  
 DE 1966-B89476 A 19661020  
 DE 1966-B90062 A 19661129

IT 24007-97-4P 24008-01-3P 24008-02-4P  
 RL: SPN (Synthetic preparation); PREP (Preparation)  
 (preparation of)  
 RN 24007-97-4 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethylethyl)amino]-1-hydroxyethyl)-3-methoxy-  
 (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

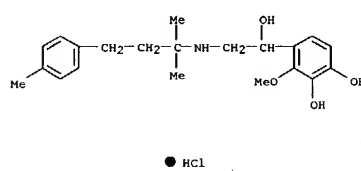
RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

L4 ANSWER 8 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)  
 $\alpha$ -[2-(methoxyphenyl)isopropylamino]-2-methyl-3,4-dimethoxyacetophenone-HCl, 205°;  $\alpha$ -[2-(p-hydroxyphenyl)-isopropylamino]-2-methyl-3,4-dihydroxyacetophenone-HBr, 115-25° (HCl salt m. 120-35°); 2-hydroxy-3,4-diphenylmethylenedioxyacetophenone, ;  $\alpha$ -bromo-2-methoxy-3,4-diphenylmethylenedioxyacetophenone, 137°;  $\alpha$ -(cyclopentylamino)-2-methoxy-3,4-dihydroxyacetophenone-HCl, 202-3°;  $\alpha$ -(benzyl-phenoxyethylamino)-2-methoxy-3,4-diphenylmethylenedioxyacetophenone-HCl, 159-61°;  $\alpha$ -(phenoxyethylamino)-2-methoxy-3,4-diphenylmethylenedioxyacetophenone-HCl, 190-2°;  $\alpha$ -(phenoxyethylamino)-2-methoxy-3,4-dihydroxyacetophenone-HCl, 174-5°;  $\alpha$ -(p-tolyloxyethylamino)-2-methoxy-3,4-dihydroxyacetophenone-HCl, 181-2°;  $\alpha$ -(tert-butyloxamino)-2-methoxy-3,4-diphenylmethylenedioxyacetophenone-HCl, 182-3°;  $\alpha$ -(tert-butyloxamino)-2-methoxy-3,4-dihydroxyacetophenone-HCl, 189-90°;  $\alpha$ -(o-tolyloxyethylamino)-2-methoxy-3,4-dihydroxyacetophenone-HCl, 197-9°;  $\alpha$ -(m-tolyloxyamino)-2-methoxy-3,4-dihydroxyacetophenone-HCl, 170-2°;  $\alpha$ -(o-methoxyphenoxyethylamino)-2-methoxy-3,4-dihydroxyacetophenone-HCl, 152-3°;  $\alpha$ -(1,1-dimethyl-3-phenylpropylamino)-2-methoxy-3,4-diphenylmethylenedioxyacetophenone-HCl, 174-6°;  $\alpha$ -(1,1-dimethyl-3-phenylpropylamino)-2-methoxy-3,4-dihydroxyacetophenone-HCl, 173-5°;  $\alpha$ -(1,1-dimethyl-3-p-tolylpropylamino)-2-methoxy-3,4-dihydroxyacetophenone-HCl, 166-7°;  $\alpha$ -bromo-2-methyl-3,4-bis(benzyloxy)acetophenone, 123°;  $\alpha$ -(tert-butyloxamino)-2-methoxy-3,4-bis(benzyloxy)acetophenone, 111-12°; 2-methoxy-3,4-bis(diphenylmethylenedioxy)butyrophenone, ;  $\alpha$ -(isopropylamino)-2-methoxy-3,4-dihydroxybutoxyphenoxy-HCl, 93-4°;  $\alpha$ -bromo-2-methoxy-3,4-diphenylmethylenedioxybutyrophenoxy, ;  $\alpha$ -(isopropylamino)-2-methoxy-3,4-diphenylmethylenedioxybutyrophenoxy-HCl, 93-4°;  $\alpha$ -(isopropylamino)-2-methoxy-3,4-dihydroxybutoxyphenoxy-HCl, 188-90° (decomp.); 1-(3,4-diaceoxy-2-methylphenyl)-1-oxo-2-isopropylaminoethane-HCl, 156°; and 1-(3,4-diaceoxy-2-methoxyphenyl)-1-oxo-2-isopropylaminoethane-HCl, 166-7°. I show sympathomimetic properties and dilate the peripheral vessels.

ACCESSION NUMBER: 1970-31420 CAPLUS  
 DOCUMENT NUMBER: 72-31420  
 TITLE: 1-(2-Substituted-3,4-dihydroxyphenyl)-2-(substituted-amino)ethanol  
 PATENT ASSIGNEE(S): Boehringer, C. H., Sohn  
 SOURCE: Fr., 12 pp.  
 CODEN: FRXXAK  
 DOCUMENT TYPE: Patent  
 LANGUAGE: French  
 FAMILY ACC. NUM. COUNT: 3  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FR 7338	M	19691013	FR 1968-7338	19680118
DE 1543372	A	19710401	DE 1966-B89417	19661018
DE 1543374	A	19720420	DE 1966-B89476	19661020
ES 346057	A1	19690316	ES 1967-346057	19671014
CH 490323	A	19700515	CH 1967-490323	19671016
AT 285582	B	19701110	AT 1967-9349	19671016
AT 288357	B	19701310	AT 1969-11652	19671016
AT 288358	B	19701310	AT 1969-11653	19671016

L4 ANSWER 8 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)  
 RN 24008-02-4 CAPLUS  
 CN 1,2-Benzenediol, 4-[2-[(1,1-dimethyl-3-(4-methylphenyl)propyl)amino]-1-hydroxyethyl]-3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-(2-[(1,1-dimethyl-3-phenylpropyl)amino]-1-hydroxyethyl)-  
 3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)

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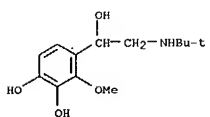
RN 24008-01-3 CAPLUS  
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L4 ANSWER 9 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN  
 GI For diagram(s), see printed CA issue.  
 AB I, having broncholytic, antipruritic, and peripheral vasodilatory  
 activities, are prepared via III and II by sequential debenzylation with  
 H over Pd on C in MeOH at 60° and 5 atmospheric and then hydrogenation over  
 Pt or Raney Ni in MeOH. Alternately for X = PhCH<sub>2</sub>, II are reduced to I  
 with NaBH<sub>4</sub> followed by debenzylation as above. III are prepared by  
 treatment of the appropriate  $\alpha$ -bromoacetophenone with R1R2NH. Of  
 protecting groups used, X and (or) X' = Me are removed by 1.5-hr. reflux  
 in 40-50% HBr and (XX' =) Ph<sub>2</sub>C (introduced by the action of Ph<sub>2</sub>CCl<sub>2</sub> and  
 pyridine in Me<sub>2</sub>CO) is removed either during the debenzylation or by 2-hr.  
 reflux in a concentrated HCl-MeOH mixture I and II prepared are  
 tabulated. Addnl.  
 described was IV.HCl (R = Pr, R<sub>1</sub> = iso-Pr, X = H, X' = Me), m.  
 93-5°. V described were (R, R<sub>1</sub>, X, X', m.p. given): Me, Ph(CH<sub>2</sub>)<sub>3</sub>,  
 Me, Me-(hydrogen oxalate m. 118-21° (Et<sub>2</sub>O); MeO, PhOCH<sub>2</sub>CH<sub>2</sub>, (XX'  
 =)Ph<sub>2</sub>C, 159-61° (CH<sub>2</sub>C<sub>12</sub>-Et<sub>2</sub>O). The preparation of several  
 intermediates is also given.

ACCESSION NUMBER: 1969:523933 CAPLUS  
 DOCUMENT NUMBER: 71:123933  
 TITLE: Broncholytic phenyl alkanolamines  
 INVENTOR(S): Mentrup, Anton; Schremm, Kurt; Thomae, Otto; Zeile, Karl  
 PATENT ASSIGNEE(S): Boehringer Ingelheim G.m.b.H.  
 SOURCE: S. African, 26 pp.  
 CODEN: SFXXAB  
 DOCUMENT TYPE: Patent  
 LANGUAGE: Russian  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

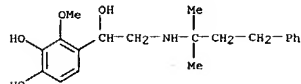
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ZA 6802425		19681108	ZA	19680417
IT 24007-97-4P	24008-01-3P	24008-02-4P		

RL: SPC (Synthetic preparation); PREP (Preparation)  
 (preparation of)  
 RN 24007-97-4 CAPLUS  
 CN 1,2-Benzenediol,  
 4-[2-[(1,1-dimethylethyl)amino]-1-hydroxyethyl]-3-methoxy-  
 (9CI) (CA INDEX NAME)



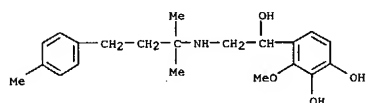
RN 24008-01-3 CAPLUS  
 CN 1,2-Benzenediol,  
 4-[2-[(1,1-dimethylethyl)amino]-1-hydroxyethyl]-3-methoxy-  
 (9CI) (CA INDEX NAME)

L4 ANSWER 9 OF 9 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)



● HCl

RN 24008-02-4 CAPLUS  
 CN 1,2-Benzenediol, 4-[2-[(1,1-dimethyl-3-(4-methylphenyl)propyl)amino]-1-hydroxyethyl]-3-methoxy-, hydrochloride (9CI) (CA INDEX NAME)



● HCl

=> logoff y		
COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	48.12	203.75
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-6.30	-6.30

STN INTERNATIONAL LOGOFF AT 13:27:55 ON 10 NOV 2004